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- Retort pouch and coextruded film therefor.
- A seal layer for use in large institutional-sized retort pouches comprises an ABA film structure wherein the A layers are each composed of a blend of a minor amount of an elastomer and a major amount of a polyclefin, and the B layer is composed of a blend of a major amount of an elastomer and a minor amount of a polyclefin. Retort pouches made with this ABA film structure as the seal layer exhibits improved impact strength.

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RETORT POUCH AND COEXTRUDED FILM THEREFOR

BACKGROUND OF THE INVENTION

This invention relates generally to retort pouches. In one aspect it relates to a coextruded three layer film for use in retort pouches wherein each layer contains elastomer. In another aspect, the layer are attent to a three layer laminate, one of which is a coextruded three ply film, for use in retortable pouches in still another aspect the invention relates to a method of packaging food for retorting.

10 Description of the Prior Art

There is a trend in the food packaging industry to replace metal cans with placific containers, notably pouches containing sterilized or pre-cooked foods. Such retort pouches offer many advantages over rigid metal packages: faster cook-sterilizing time, less shelf storage space, easier disposal, improved food taste, etc. The pouches are typically three layer laminations of an outer film (e.g., biax oriented hylon and/or polyester), a middle moisture and oxygen barrier film (e.g., aluminum foil, PVDC, or EVOH), and an inner heat seal layer film (PP, PPRC, or LLDPE or blends thereof). Adhesives to join the three layers together are typically of the polyurethane and/or maleated polyolefin types. (The resin abbreviations used herein are defined in "Description of Preferred Embodiments".)

Large No. 10 cans (those containing about 3.2 liters of liquid food) have been flow to convert to flexible retort pouches. Large plastic containers used to package foods have been limited to high acid foods (e.g., catsup) which do not require sterilization to kill botulism toxin. In such containers, LLOPE is often selected as the inner heat seal layer film because of its low cost, toughness and drop impact resistance. However, LLDPE cannot be used in large institutional-size retort pouches (volumetric content larger than liter) which require sterilization, because it softens at typical high temperature sterilization or retort temperatures (121°C and higher). Although PP or PPRC films can be and are used in smaller size retort pouches (in the order of 0.25 to 1 liter content) because of their higher temperature resistance, they are unacceptable in the larger size pouches because they do not possess adequate toughness or drop impact strength to prevent breakage. With small containers, hydraulic forces resulting from vertical impact are not large. However, with large containers, the hydraulic forces become excessive, particularly on lateral seats. Because of this deficiency there exists a definite market need for institutional size pouches (greater than 1 liter and preferably 2 liter and larger) possessing a tough inner ply heat seal film which is resistant to high temperature retorting temperatures.

Blending elastomer into PP and PPRC is one well established method for improving film toughness and is being practiced commercially. PPRC monolayer film modified with PIB is used as the inner seal layer of medical enteral pouches used to deliver nutritional fluid to hospital patients unable to take food erally. These small 3-layer laminated pouches are typically one liter in size and weigh approximately 2.2 lbs. At least one side of these pouches consists of an all-plastic construction so that the fluid level can be monitored. In order to assure adequate impact toughness, these pouches are subjected to vertical drop tests. Typical vertical drop heights for these products are 4-6 feet. For greater drop height corromance, smaller pouches (in the order of 0.5 liters) are sometimes used. However, pouches of this type are far too small for use as institutional food pouches.

European Patent Application 0.165,791 discloses a composite film structure comprising rubber-containing core and polyolefin resin. The skin layers of this composite however, contain no rubber. As demonstrated below, rubber (elastomer) in both the skin and core layers improves performance of the pouch

Summary of the Invention

It has been discovered that by utilizing a special coextruded inner heat seal layer in a retortable pouch, the problem of room temperature impact strength of large containers is effectively overcome. Although retortable pouches are available in a variety of sizes and constructions, those used in food backaging almost all include a barrier layer and an inner seal layer. The inner seal layer of the retortable bouch of the present invention comprises a coextruded multilayer film which may be an ABA or an ABC stricture, where A is the inner skin layer. Of the outer layer, and B the core layer. As used herein, the inner layer as is the

layer of the package in contact with the food and the outer layer C is adjacent to the barr or layer with the without adhesives).

The A and B layers comprise a blend of an elastomer and a polyolefin, preferably PP, more preferably PPRC. Outer skin layer C may be any polyolefin suitable for interfacing with the barrier resinfol, our preferably has the same composition as A, making the seal layer an ABA structure. Layers A and B each include sufficient elastomer to increase impact strength to the final structure. However, skin layer A contains substantially less elastomer than core layer B. The amount of elastomer in the total film should be in excess of 50 wt% but not more than 70 wt% and preferably not more than 60 wt%. The amount of elastomer in skin layer(s) A should be sufficiently small to avoid or minimize blocking or sticking on the film is 1 aut sufficiently large to add to the impact strength of the 3 layer structure. ABA ABC thickness ratios may vary within wide limits but preferably range from about 3:94:3 to 20:60:20.

The present invention also includes a pouch structure per se, comprising an outer structure! laver a barrier layer and the three ply inner seal layer of the structure described above. The three ply ABA or ABC structures are useful in retortable pouches which exhibit exceptionally high impact strength and are particularly suitable for use in large institutional-sized retortable flexible containers darger than 1 liter e.g., 1.25 liters and preferably between 1.5 and 4.0 liters and most preferably between 1.8 and 3.5 liters larger.

Description of the Drawings

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Figure 1 is a cross section view of a laminated film constructed according to the cresent invention. Figure 2 is a pouch made from the film of Figure 1.

25 Description of the Preferred Embodiments

In its broadest embodiment, the invention comprises a multilayer film which is particularly users, in retort pouches. In many institutions such as hospitals, military, and schools, large retort pouches have potential application. A retort pouch must possess the following properties, all of which are provided or affected by the inner seal layers:

- 1. they must be heat sealable for closing the container securely following filling:
- 2. they must be flexible and yet have sufficient toughness and impact strength to enable the pouch to withstand severe abuse testing in accordance with industry shipping standards.
- 3. they must be capable of withstanding sterilization at temperatures in the range of 121°C 135°C to kill botulism:
- 4, they must possess non-blocking properties in that the interior opposed faces of the container must not stick together which would impede filling of the pouch:
- 5. the film used in the pouches must be processable without sticking as it is unrolled from the film roll during manufacture of the pouches and subsequent sterilization:
- 6. they must have sufficient capacity to be economical. (Institutions serving large numbers of people are not likely to use extensively containers having capacities less than one liter.)

Although retortable pouches, one panel of which is represented as 10 in Figure 1, are available in various constructions, they generally include the following layers: an outer structural layer 11, a barrier layer 12, and a seal layer 13 laminated together with suitable adhesive tie layers. The retort pouch layers are described separately below.

Structural Layer: The structural layer 11 provides structural strength, abrasive resistance and a printable surface for the pouch. It normally is composed of a thermoplastic such as polyolefins, polypropylene [e.g., propylene homo polymers (PP) and copolymers], polyester, nylon, and the like. Biaxially oriented hylon and biaxially oriented polyester are the preferred structural layers. Although not essential to the pouch, the structural layer is preferred since as noted above it adds strength, durability, and printability to the pouch.

Barrier Layer: The barrier layer 12 is substantially impermeable to oxygen for protecting the food medium contained in the pouch. Barrier layer materials include aluminum foil, thermoplastic films such as ethylene vinyl alcohol (EVOH), polyvinylidenedichloride (PVDC), polyvinyl alcohol(PVA), and the ike. Aluminum foil is the preferred barrier layer. In packaging where oxygen barrier requirements are not stringent, biaxially oriented hylon or polyester are sometimes employed. The compositions and methods of manufacture of barrier layers and structural layers are well known to the industry and are described in the published literature.

Seal Layer: The main function of the seal layer 13 is to provide a substance which is hoat scalable

(fusible) with itself to seal the pouch.

An important aspect of the present invention is the composition of the seal layer which unables that pouch to satisfy the six requirements mentioned above. The present invention contemplates the use of a coextruded film as the seal layer, comprising an ABA composite or an ABC composite wherein A is the inner layer of the film. B is a core layer, and C is the outer layer of the film.

The inner layer (A) [and preferably the outer layer (C)] and the core layer (B) each comprises this essential ingredients: a polyolefin and an olefinic elastomer which are blended in different races. The core layer is elastomer rich - more than 50 wt% elastomer - and the inner layer (and preferably the latter layer) is elastomer lean - less than 50 wt% elastomer.

The polyolefin resins which are suitable for use in the practice of this invention for bonding with the elastomer are those which have a semi-crystalline or crystalline meiting point of at least 11030 illustrative. non-limiting examples of the polyolefins suitable for use in the practice of this invention are convercelylene (PP), linear low density polyethylene (LLDPE) (e.g., a copolymer of ethylene and 3-15% or so elefth having 4 to 12 carbon atoms) and polybutylene (PB). As used in the specification and claims, the term "polypropylene" includes homopolymers of propylene as well as reactor socialymers of cropylene PPRC: which include both random copolymers and impact copolymers. Random popolymaks contain about 1 to about 6 wt% ethylene or an alpha olefin monomer of 4 to 16 carbon atoms, and impact cooclymers contain from 6 to 20 wt% ethylene or alpha olefin comonomer of 4 to 16 carbon atoms. The polypropylene can be highly crystalline isotactic or syndiotactic polypropylene. The density of the PP or PPRC ranges from about 0.89 to about 0.91g.cc and the MFR is typically in the range of 0.5 to 5.0. Blends of these polyolefin resins may be used. For example, LLDPE may be blended into PP, not at a high enough concentration to harm thermal stability, but at a concentration to favorably affect heat seatability. The propylere content of the inner layer A should be at least 20 wt%.

As demonstrated by the experiments described below, PPRC (specifically random copolymer) gave the best results in improvements in the impact strength and accordingly is preferred for blending with the elastomer resins.

The rubbers (elastomeric resins) which can be used in the practice of this inventic, include both synthetic and natural rubbers; preferably the rubbers have a glass transition temperature (Tg) of less than 0°C. Illustrative, non-limiting examples of rubbers suitable for use in the practice of this invention include polyisobutylene (PIB), butyl rubber, halogenated butyl rubber, ethylene propylene rubber (EPM), ethylenepropylendiene rubber (EPDM), polyisoprene, polychloroprene, styrene-butadiene rubber, polyoutene copolyers, nitrile rubbers, chlorosulfonated polyethylene, etc. while polyisobutylene (PIB) is not a true rubber because it cannot be vulcanized, it can be utilized in the practice of this invention provided that the PIB has a viscosity average molecular weight (Flory) of about 900,000 to about 1.6 million.

The term "rubber" as used in the specification and claims means any natural or synthetic colymer which can be vulcanized or cured so as to exhibit elastomeric properties. For the purpose of this invention, PIB is considered a rubber although it cannot be vulcanized.

The terms EPM and EPDM are used in the sense of their ASTM designations (ASTMD-1418-72a). EPM is an ethylene-propylene copolymer which can be cross-linked by radiation curing or peroxide curing

As used in the specification and claims the term "EPDM" means terpolymers of ethylane and aiphaolefin and non-conjugated diene. The non-conjugated diolefin can be straight chain, branched chain or cyclic hydrocarbon di-olefins having about 6 to about 15 carbon atoms such as:

- A. straight chain dienes such as 1.4-hexadiene and 1.6-octadiene:
- B. branched chain acyclic dienes such as 5-methyl-1, 4-hexadiene: 3.7-dimethyl1-1,6-octadiene: 3.7dimethyl1-1, 7-octadiene and the mixed isomers of dihydro-myricene and dihydrocinene:
 - C. single ring alicyclic dienes such as 1,3-cyclopentadiene: 1,4-cyclohexadicne: 1,3-cyclopentadiene and 1.5-cyclododecadiene:
- D. multi-ring alicyclic fused and bridged ring dienes such as tarahydroindene, matnyl, dicyclopantadiene: bicyclo-(2.2.1)-hepta-2, 5-diene: alkenyl, alkylidene, cycloalkenyl and cycloalkylidene norbornanes 50 such as 5-methylene-2-norbornene (MNB), 5-propenyl-2-norbornene, 5-isopropylidene-2-norbornene, 5-idcyclopentenyl)-2-norbornene. 5-cyclohexylidene-2-norbornene. 5-vinyi-norbornene and norcornadiane

Of the non-conjugated dienes typically used to prepare EPDM teruplymens the proferred cients are dicyclopentadiene, 1-,4-hexadiene, 5-methyl-2-norbornene and 5-ethylicisne-2-norbornene Particularly preferred diolefins are 5-ethylidene-2-norbornene (ENB) and 1.4-hexadione.

EPDM elastomers and their general method of manufacture are while known in the art. The professed EPDM elastomers contain about 20 to about 90 wt% ethylene, more preferable strong 30 to 30 wt%; ethylene, most preferably about 35 to about 75 wt% ethylene.

The alpha-olefins suitable for use in the preparation of EPOM are professibly CHO uses a forest

Illustrative non-limiting examples of such alpha olefins are propylene. 1-butene. 1-centere. 1-herzne. 1-bettere octene and 1-dodecene. The alpha olefin is generally incorporated into the EPDM polymer at about 10 to about 80 wt%, more preferably at about 20 to about 70 wt%. The non-conjugated diene is incorporated into the EPDM at about 0.5 to about 15 wt%; more preferably about 1 to about 5 wt%, e.g., 3 wt%.

The term "nitrile rubber" means an acrylenitrile copolymer rubber. Suitable nitrile rubbers comprise rubbery polymers of 1.3-butadiene and about 20-50 wt% acrylenitrile. Any nitrile rubber which is a "solid" rubber having an average molecular weight of at least 50,000 and preferably between about 100,000-1,000,000 can be used. Commercially available nitrile rubbers suitable for the practice of the invention are described in Rubber World Blue Book. 1980 Edition, "Materials and Compounding Ingredients for Publication," pages 386-406, which is incorporated herein by reference.

Butyl rubber is a copolymer of an isoolefin and a conjugated multiolefin. The useful roctiveners comprise a major portion of isoolefin and a minor amount, preferably not more than 30 with, of a conjugated multiolefin. The preferred copolymers comprise about 85-99.5 wt. (preferably 95-99.5 wt%) of a Ci-Cr isoolefin, such as isobutylene, and about 15-0.5 wt% (preferably about 5-0.5 wt%) of a multiolofin of about 4-14 carbon atoms. These copolymers are referred to in the patents and literature as "butyl rubber"; see, for example, the textbook Synthetic Rubber by G. S. Whitby (1954 edition by John Wriey and Sons, Inc.), pages 608-609, etc. which is incorporated herein by reference. The term "butyl rubber" as used in the specification and claims includes the aforementioned copolymers of an iscolefin having 4-7 carbon atoms and about 0.5 to 20 wt% of a conjugated multiolefin of about 4-10 carbon atoms. Preferably these copolymers contain about 0.5 to about 5% conjugated multiolefin. The preferred isoolefin is isobutylene. Suitable conjugated multiolefins include isoprene, butadiene, dimethyl butadiene, piperylene, etc.

Commercial butyl rubber is a copolymer of isobutylene and minor amounts of isoprone. It is generally prepared in a sturry process using methyl chloride as a vehicle and a Friedel-Crafts catalyst as the polymerization initiator. The methyl chloride offers the advantage that AICI₃, a relatively inexpensive Friedel-Crafts catalyst is soluble in it, as are the isobutylene and isoprene componemers. Additionally, the butyl rubber polymer is insoluble in the methyl chloride and precipitates out of solution as fine partic. The polymerization is generally carried out at temperatures of about -90°C to -100°C. See U.S. Patent Nos. 2.356.128 and 2.356.129 incorporated herein by reference.

In the halogenation process, butyl rubber in solution is contacted with chlorine or bromine in a series of high-intensity mixing stages. Hydrochloric or hydrobromic acid is generated during the halogenation step and must be neutralized. For a detailed description of the halogenation process see U.S. Patent Nos. 3.029.191 and 2.940,960, as well as U.S. Patent No. 3.099,644 which describes a continuous chlorination process, all of which patents are incorporated herein by reference.

The rubbers used in the practice of this invention are preferably utilized in their unvulcanized state. The preferred rubber is PIB.

The core layer (B) and skin layers (A and C) of film composite 13 preferably have the following construction:

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	•	Compositi	on (wt%)	Thickness
		Elastomer	Polyplefin	(% of Total)
	Outer Layer (C)	0-40	60-100	3-20
5	Core Layer (B)	>50-75	25-<50	50-94
	Inner Layer (A)	10-49	60-90	3-20
o	Total Composite	> 50-70	< 30-50	2-10 mis
		Preferred C	onstruction	
5				Thislenges
		Composit	ion (wt%)	Thickness
		Elastomer	Polyolefin	(∦ of Total)
c	Outer Layer (C)	15-30	70-85	7-15
•	Core Layer (B)	>55-70	30-45	70-86
	Inner Layer (A)	15-30	70-35	7-15
25	Total Composite	> 50-60	40-<50	4-7 mi]

As indicated above the total elastomer content of the composite seal layer 13 comprises more than 50% in order to impart the necessary impact strength to the pouch. The polyoletin, particularly PPRC, provides for shrinkage stability of the film and retortability without sacrificing impact strength. The elastomer in the inner layer A contributes to impact strength and heat seal strength; excessive elastomer therein is undesirable since it results in roll sticking and/or container blocking. The gradient approach (i.e., rubber lean skin and rubber rich core) yields the highest total rubber content without film blocking.

The compositions of this invention may also include rubber extender oils as well as processing aids and stabilizers, fillers and the like.

Food pouches are generally steam autoclaved at about 121°C in order to sterilize the food. The pouches prepared with the use of composite film as the seal layer 13 according to this evention must not only withstand the autoclave conditions, they must also exhibit a low water loss both during autoclaving and in storage. Rubbers based on isobutylene, e.g., PIB and butyl rubber, have the lowest water vapor transmission rate among rubbers, and also are broadly FDA regulated. Hence, they are the preferred rubbers for use in this invention.

Preparation of compositions for each layer usable in this invention can be achieved in several different ways. The components may be brought into intimate contact by, for example, dry blending these materials and then passing the overall composition through a compounding extruder. Alternatively, the components may be fed directly to a mixing device such as a compounding extruder, high shear continuous mixer, two roll mill or an internal mixer such as a Banbury mixer. The optional ingredients previously described can be added to the composition during this mixing operation. It is also possible to achieve melt mixing in an extruder section of a coextrusion apparatus. Overall, the objective is to obtain a uniform dispersion of all ingredients and this is readily achieved by inducing sufficient shear and heat to cause the plasues component(s) to melt. However, time and temperature of mixing should be controlled as is normally done by one skilled in the art so as to avoid molecular weight degradation.

Although ABC structures may be employed for fabrication by coextrusion, as mentioned previously, it is preferred to use an ABA film. This film may be coextruded by using conventional extruding and casting equipment wherein one extruder casts the core layer (B) and another extruder casts the skin layers (A).

Chill rolls and silicon rolls also may be employed in the casting process. The three layer coex film may be prepared and stored as a roll for fabrication into the pouch film in subsequent operations.

RETORT POUCH

As mentioned previously, the composite film forms the inner seal layer of a retort pouch 10A having the following structure:

•	•			THICKNESS		
						Preferred
			1	Broad Range		Range
10		COMPOSITION	mils	% of Total	<u>mils</u>	% of Total 5-20
	Structural Layer	tnermo- plastic		0-25		O⇔47/
:5	Barrier Layer	barrier		3-15		4-10
29	Seal Layer	resin/foil composite film	2-10	50-97	4-7	70-90

As shown in Figure 2, the pouch 10A preferably is made by laminating the three layers 11, 12, and 13 forming a roll stock having the cross section of Figure 1. The roll stock then is used to form two panels 14 and 15, cutting the panels to the proper size, and heat sealing three of the four sides of the aligned edges 16, 17 and 18 of the panels, leaving one side open 19 to permit filling with food or other perishable goods.

The three layers may be laminated together using conventional equipment and techniques.

Suitable adhesives, or tie layers, may be used to adhere the component layers together. For example, adhesives such as heat curable polyurethane are useful at the interface of hylon layer 11 and aluminum foil layer 12 and at the interface of aluminum layer 12 and the seal layer 13. Likewise, maleic-grafted polyolefins may be used at the interface of nylon layer 11 and PVDC or EVOH 12, and at the interface of the ABA (or ABC) seal layer 13 of the present invention. Other tie resins are well known in the art.

The laminated roll stock is processed into pouches using conventional equipment and techniques. The form, fill and seal (F.F.S) machinery process is one such technique. In this process, two superimposed layers of the laminated film of the desired width with interfacing seal layers are fed into the packaging machine. Bar seals fuse three sides of the square or rectangular pouch and a jaw separates the pouch with one open end. Food is then introduced into the pouch through the open end. Finally, the machine heat seals the open end producing a fully closed pouch.

For large institutional containers, the panels for pouches will be sufficiently large to provide the peuch with a capacity ranging from greater than 1 (e.g., about 1.2 liters) to 5 liters, preferably between 1.5 to 4.0 liters, and most preferably between 1.8 and 3.5 liters. These pouches are adapted to be filled with a liquid semiliquid food (e.g., low acid particulates such as stew). Packaging machines which fill by the FFS process include Prodo Pak's Model HFF-1000 and N & W Packaging System Model MAX 2014.

Preformed pouches may also be employed in this operation, pouches with an open end are precared and subsequently filled with the liquid or semiliquid substance in a separate step.

The packaged food is then sterilized at temperatures of 121°C to 135°C for 3 to 60 minutes depending on the type of food and food processing techniques.

One of the problems associated with institutional-sized plastic containers for packaging liquid or semiliquid food is their inability to withstand impact forces following heat sterilization. The heat weakens the seals and the hydraulic forces on dropping creates high stresses in the horizontal direction. Thus, an essential property of each pouch is that it is capable of withstanding impacts during handling and transport. The industry has developed a number of tests for determining the suitability of the pouches for shipping in cases; two of which are ASTM and NSTA standards described below. In addition, it has been found that drop tests on individual pouches provides a reliable test for determining (qualitatively) the impact strength of pouches. These tests involve multiple drops of each pouch before retorting. The term "multiple drop tests" means that the pouch is dropped on each of the four sealed edges and on one flat surface from an clovation of 10 feet onto a flat metal surface.

EXPERIMENTS

Effects of Rubber on Heat Sealability of PP:

Tests were carried out on the heat seal strength of monolayer film of various blends of PP and EPDM (Vistalon 93708 marketed by Exxon Chemical Company). These tests indicated that the addition of the elastomer up to about 40 wt% improves heat seal strength for elastomer PP blend film. Elastomer content above 40% in the film showed little or no improvement in heat seal strength. Reterting a blend of 40% elastomer and 60% PP at 121°C for 30 minutes reduced seal strength by only 19%; the remaining seal strength was adequate for packaging purposes.

Pouch Tests

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Pouches (12 inch * 12 inch and 10 inch * 13 inch) were made using a 4-mil or 7-mil seaf layers having the following compositions:

TABLE 1 - SEAL LAYER FILM X

20	Layer	Composition	wts	Thickness (% of Total)
	Δ	VISTANEX L-1001/PPRC2	24/75	10
	8	VISTANEX L-1001/PPRC2	65/35	<i>2</i> 0
<i>2</i> 5	C	VISTANEX L-1001/PPRC2	24/75	10

1 pI3, EXXON Chemical Co.
2 3 wt% etnylene, 1 MFR

The seal film was laminated with structural and barrier layers to form two roll stocks which were used to labricate pouches having the compositions of Pouches 1 and 2 described in Table 2. The Pouch 2 structures made from one of the roll stocks were 12" 12" and were fabricated, filled, and sealed using Prodo Pak's Model HFF1000 Filler/sealer. The seal temperatures were about 410°F with a horizontal dwelf of 4 seconds and a vertical dwell of three-fourths second and a line speed of 12 pouches per minute. Seal pressures were 80 psi for horizontal seals and 70 psi for vertical seals. Pouch 1 structures made from the other roll stock were 10" 13" and were made using preformed equipment at approximately the same process conditions.

TABLE 2 - LAMINATED POUCHES

	Pouch 1	Pouch 2
Structural Layer:	PET/T3	BN/T3
Barrier Layer:	A1/T4	A1/T4
Seal Layer:	Seal Layer X	Seal Layer X

BN - Biax nylon (60 gauge)

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Al - Aluminum Foil (35 gauge)

T₃ - Tie layer marketed by Morton Chemical as Adcote 550.

T₄ - Tie layer marketed by Morton Chemical as Adcote 506-40/9L10.

PET - Biaxially oriented polyester manufactured by DuPont (48 gauge).

The pouches were filled with 1.9 liters or 3.18 liters of liquid, and drop tested by the multiple drop test method before retorting. The following summarizes the test results:

TABLE 3 - MULTIPLE DROP TESTS

25		Pouch 1 (1.9 Liters)	Pouch 2 (3.18 Liter)
	No. Samples Tested	50 (4 mil)	75 (4 mil)
	(Thickness of Seal Layer)		50 (7 mil)
30	Results	Passed multiple	Passed multiple
	230103	drop test	drop test

The above tests demonstrate that (a) the retorted pouches made in accordance with the present invention exhibit improved impact strength and (b) the improved results are due to the composition of the seal layer.

In order to assure safe handling of the pouches packed in shipping containers during transit, the pouches (after filling and retorting) should pass tests established by the ASTM and/or the NSTA. Such test include Vibration Testing for Shipping Containers (ASTM D999-75) and Drop Testing for Shipping Containers (ASTM D775-61), and NSTA Test Procedures, Project IA. The pouches of the present invention packed in shipping containers (2 stacks, 3 high) are capable of passing the ASTM tests and the more stringent NSTA tests. This capability of the institution sized packages, particularly those between 1.3 and 3.5 liters has not been achieved with prior art pouches.

The following Container Drop Tests on Pouches 1 and 2 demonstrate this capability and exemplify the high impact strength of the pouches.

Pouches had the structures and compositions of Pouches 1 and 2 described in Table 2 and were retorted for about 60 minutes at about 121°C. Corrugated cardboard cases were packed with six of each type of pouch after retorting. Each case was provided with two vertical compartments which snugly contained three stacked pouches. Each case was tested by the ASTM or NSTA Drop Tests described above (including vibration tests prior to dropping). TABLE 4 summarizes the Container Drop Tests.

TABLE 4 - CONTAINER DROP TESTS

		Seaī					
5		Layer			No.		
		Thick-		Liquid	Con-		
	Pouch	ness	size	Contents	tainers	Test	ïest
	Type	Mils	inches	liters	Tested	<u>Method</u>	Results
•0	1	4	10·x 13	about 1.9	20	ASTM 0775-61 (12")	Passed
	2	4	12 x 12	ahout 3.1	S 10	ASTM D775-61 (12")	
	_					and NSTA	Passed
15	2	7	12 x 12	about 3.1	.S 10	ASTM 0775-61	
	_	·				and NSTA	Passed
	1	7	10 x 13	about 1.9	9 . 5	ASTM 0775-61 (12)	Passed

As indicated above the Multiple Drop Tests are carried out before retorting and the Container Drop Tests are carried out after retorting. Tests have shown that pouches that pass the Multiple Drop Tests also pass the ASTM and NSTA Container Drop Tests. The Multiple Drop Test thus provides a reliable, simple and inexpensive test for determining the quality of shipping pouches.

Blocking Tests

Tests using a roll of monolayer seal layer film containing 65 wt% VISTANEX and 35 wt% PPRC exhibited severe blocking, demonstrating the problem associated with high rubber content. The roll stock used in preparing Pouches 1 and 2, however, exhibited little or no blocking.

Claims

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- 1. Multilayer heat sealable and retortable film which comprises:
- (a) an inner layer comprising a blend of a major weight portion of a polyolefin and a minor weight portion of an elastomer, said inner layer being heat sealable with itself.
- (b) a core layer comprising a blend of a major weight portion of an elastomer and a minor weight portion of a polyolefin resin; and
- (c) an outer layer of a polyolefin; the total elastomer content in the layers is more than 50 wt% thereof.
- 2. Multilayer film according to claim 1 wherein the polyolefin in the core layer and or inner 'ayer is a propylene polymer, preferably PPRC such as propylene impact copolymer centaining from 6 to 20% of ethylene.
- 3. Multilayer film according to claim 1 or claim 2 wherein the elastomer in the inner and or core layer is PIB.
- 4. Multilayer film according to any of the preceding claims wherein the said outer layer comprises from 0 to 40 wt% of an elastomer and from 60 to 100 wt% of a polyolefin; the core layer comprises more than 50 wt% and not more than 75 wt% of said elastomer; and the inner layer comprises from 10 to 40 wt% of said elastomer, preferably with said inner layer comprising from 15 to 30 wt% of said elastomer and said core layer comprising from 55 to to 70 wt% of said elastomer.
 - 5. Coextruded multilayer film comprising an ABC layer construction wherein:
- (a) the A layer is the inner layer and comprises a blend of a propylene polymer and from 10 to 40 wt% of an elastomer, said A layer being heat sealable with itself:
- (b) the B layer is the core layer comprising from more than 50 to 75 wt% of an elastomer and from 25 to less than 50 wt% of a propylene polymer; and

- (c) the C layer is the outer layer comprising a propylene polymer and from 0 to 49 with at an elastomer, the ratio of the thickness of ABC layers being from 3:94:3 to 20:60:20; and
 - (d) the total elastomer content of layers A. B. and C is more than 50 wt% but not more than 75 Atha
- 6. Film according to claim 5 wherein the elastomer in the ABC layers is the same and is PIB and the propylene polymer in the ABC layers is the same and is PPRC such as propylene impact copic ymer containing from 6 to 20% of ethylene; the C layer is the same composition as A layer; and or wherein the elastomer concentration in layer B comprises from 55 to 70 wt% of the blend and the concentration of the elastomer in layers A and C comprise from 15 to 30 wt% thereof, preferably with the total elastomer common of the film being greater than 50 and not more than 60 wt%.
 - 7. Coextruded multilayer film useful in retortable pouches which comprises:
- (a) an inner seal layer comprising a blend of from 70 to 85 wt% of a propylene random less tymes and 15 to 30 wt% of PIB, said inner seal layer being heat fusible with itself:
- (b) a core layer comprising a blend of from 30 to 45 wt% of a propylene random copolymer and from 55 to 70 wt% of PIB:
- (c) an outer layer comprising a blend of from 70 to 85 wt% of a propylene random concurred and from 15 to 30 wt% of PiB:
- (d) wherein the thickness ratio of the three layers ranges from 3:94:3 to 20:60:20 and the elastomer content of the three layers constitutes more than 50 but not more than 60 wt%.
- 8. A film laminate comprising a film according to any of the preceding claims and a barrier layer of a as film or foil laminated to the C or outer layer of the film, preferably with the barrier layer being aluminum foil and the laminate further comprising a structural layer laminated to the barrier layer.
 - 9. Retortable pouch comprising first and second film panels, each panel comprising a film according to claim 8, said film panels having aligned edge portions heat fused together thereby defining a container having a volume capacity of more than 1 liter and not more than 5 liters.
- 10. Pouch according to claim 9 wherein the panels each include a structural layer laminated to the barrier layer and are of sufficient area to provide the container with a capacity of between 1.5 and 4.7 liters of liquid or semiliquids, and the pouch is capable of withstanding multiple drop tests when filled with a liquid or semiliquid before retorting, preferably with the pouch having a capacity of between 1.3 and 3.5 inters and, with other substantially identical pouches, being capable of withstanding case drep tests as defined in ASTM D775-61, after retorting at a temperature of at least 121°C and or the barrier layer being composed of aluminum foil and the structural layer being composed of biaxially oriented hylon or polyester.
 - 11. A retortable, heat sealed pouch comprising:

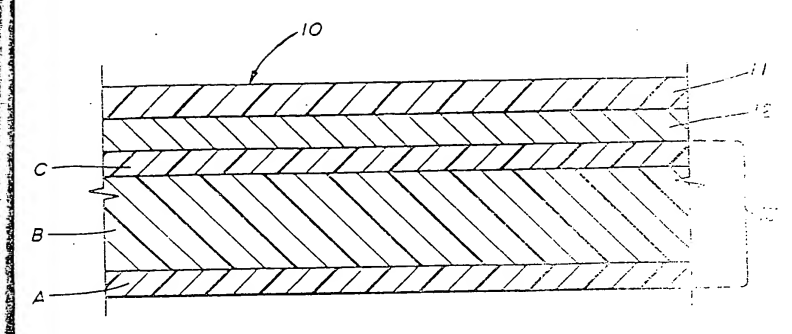
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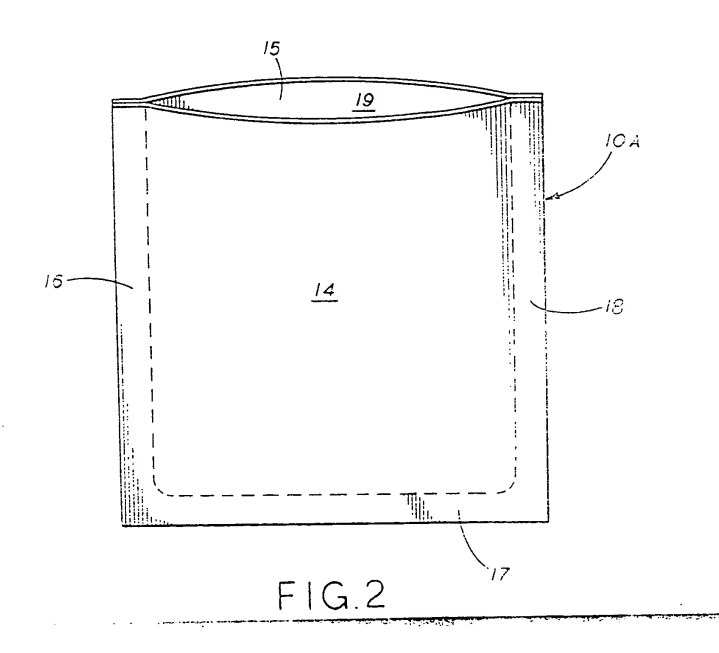
- (a) first and second panels, each comprising;
 - 6) a structural film which comprises not more than 20% of the panel thickness;
- (ii) an oxygen barrier of aluminum foil and comprising not more than 20% of the panel thickness laminated to the structural film;
- (iii) a film having the composition according to any of claims 1 to 7 and being 2 to 10 mils thick wherein the C layer is laminated to the barrier film or foil;
- (b) said panels having their edge portions heat sealed thereby defining an enclosed container having a capacity in excess of 1 liter;
- (c) a figuid or semiliquid substance contained in said enclosed container, said peuch being capable of withstanding case drop tests in accordance with NSTA Test procedures after retorting at a temperature of at least 121°C, preferably with the structural layer being biaxially oriented hylon and the pouch having a capacity of between 1.8 and 3.5 liters.
 - 12. A method of packaging liquid or semiliquid food which comprises:
- (a) forming substantially identical panels, each comprising an outer structural layer composed of a thermoplastic resin, a core layer of a barrier film or foil and a inner seal layer of extruded film according to any of claims 1 to 7:
 - (b) superimposing one panel over the other panel with interfacing inner seal layers:
- (c) heat sealing all but one edge of the aligned edges of the panels forming an interior pouch having eseparable internal panel portions and an opening at one edge portion;
- (d) filling the interior of the panels through said opening with more than 1 liter of a perishable liquid or samiliquid feed:
 - (e) heat sealing said opening thereby enclosing said food, and:
 - (f) retorting said filled pouch at a temperature about 121°C to kill botulism texin in the local and
 - (g) backaging the pouch in a case; and optionally

(h) drop testing said case in accordance with ASTM D775-61, preferably with the pouch being substantially square or rectangular in shape and area of each panel being sufficient to provide the occup with a capacity of between 1.8 and 3.5 liters.

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FIGIT



. 4. 4



EUROPEAN SEARCH REPORT

EP 88 30 0091

	DOCUMENTS CONSU	DERED TO BE RELEVAN	T		
Category	Citation of document with income of relevant pas	dication, where appropriate.	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.4)	
Y	GB-A-1 526 722 (OLE * Page 11, lines 94- 6-11; claims 1,40,41 lines 33-50 *	·103; page 11, lines	1,2,3	B 32 B 25/08 B 65 D 81/34	
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А	GB-A- 871 959 (MON * Claims 1,4,6 *	ITECATINI S.A.)	1		
				TECHNICAL FIELDS SEARCHED (Int. Cl.4)	
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-	The present search report has be	en drawn up for all claims			
	Place of search	Date of completion of the search	<u>' </u>	Examiner	
TH	E HAGUE	14-04-1988	IBA	RROLA TORRES O.M.	
i : pa do	CATEGORY OF CITED DOCUMEN rticularly relevant if taken alone rticularly relevant if combined with ano cument of the same category	E: earlier patent de after the filing sther D: document cited L: document cited	ocument, but pub date in the application for other reasons	lished on, or	
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